Lineage-Independent Tumors in Bilateral Neuroblastoma


SUMMARY

Childhood tumors that occur synchronously in different anatomical sites usually represent metastatic disease. However, such tumors can be independent neoplasms. We investigated whether cases of bilateral neuroblastoma represented independent tumors in two children with pathogenic germline mutations by genotyping somatic mutations shared between tumors and blood. Our results suggested that in both children, the lineages that had given rise to the tumors had segregated within the first cell divisions of the zygote, without being preceded by a common premalignant clone. In one patient, the tumors had parallel evolution, including distinct second hits in SMARCA4, a putative predisposition gene for neuroblastoma. These findings portray cases of bilateral neuroblastoma as having independent lesions mediated by a germline predisposition. (Funded by Children with Cancer UK and Wellcome.)
plied widely to the study of the evolution of human cancers. We have recently adapted this approach to enable the reconstruction of the embryologic features of childhood tumors. As cells divide and develop, they acquire (post-zygotic) somatic genetic variants that represent embryologic lineages and may be measurable in normal tissues. These variants and lineages render the organism mosaic. Therefore, in principle, it would be possible to relate tumors to embryologic lineages through mosaic variants (Fig. 1), which may unambiguously resolve whether tumors are developmentally related and reveal a precise developmental stage at which tumors formed or segregated. For example, in the case of bilateral Wilms’ tumor, a childhood kidney cancer, we found that tumors originated from a common embryologic precursor that formed after blood and kidney lineages segregated. Here, we applied our approach to rare cases of bilateral neuroblastoma in two children to investigate whether these lesions represented independent cancers.

**Figure 1. Possible Developmental Origins of Multisite Tumor Formation.**

The expected pattern of mutation sharing between tumors and paired normal blood reflects the origin of bilateral tumors. Bilateral tumors can arise from a common progenitor in development, which will be delineated by a shared trunk of mutations that is absent from blood (Panel A). If the tumors diverged early in embryogenesis, one or both lesions will share mutations with blood but not with the other lesion (Panel B). Letters in the phylogenetic trees refer to postzygotic mutations; the variant allele frequency of these mutations is provided in the heat maps.

**CASE REPORTS**

Patient 1 (case number, PD36812), a previously healthy 4-year-old boy, was referred to us for the evaluation of abdominal masses (Fig. 2A). After performing standard diagnostic and staging investigations that included bilateral biopsies of suprarenal lesions, we derived a diagnosis of bilateral neuroblastoma, with each tumor having poorly differentiated histologic features, no amplification in MYCN (a member of the Myc family of oncogenes), and the deletion of 19p. There was no evidence of distant metastatic disease. We initiated treatment according to the high-risk protocol of Société Internationale d’Oncologie Pédiatrique (SIOP).

Patient 2 (case number, PD34954), a 4-year-old girl, was referred to us for the management of bilateral adrenal masses that were shown to be undifferentiated neuroblastoma with widespread metastatic disease. In contrast to Patient 1, Patient 2 had a left adrenal tumor that was locally extensively invasive, whereas the right adrenal
The new england journal of medicine

November 5, 2020

1862

n engl j med 383;19 nejm.org

The New England Journal of Medicine

Downloaded from nejm.org at UNIVERSITY COLLEGE LONDON on November 25, 2020. For personal use only. No other uses without permission.
Copyright © 2020 Massachusetts Medical Society. All rights reserved.

tumor constituted a relatively small adrenal mass (Fig. 3A). A diagnosis of congenital central hypoventilation syndrome was made in association with a germline mutation in \textit{PHOX2B}, an alteration that also results in a predisposition to neuroblastoma.\textsuperscript{6}

### Genomic Analysis and Results

We performed whole-genome sequencing of DNA derived from each tumor and from peripheral blood samples obtained from each of the children and their parents. Using a validated variant-calling pipeline,\textsuperscript{5} we determined variants of all classes, which were subdivided into inherited germline, de novo germline, mosaic (embryonic), and somatic variants. From these catalogues, we reconstructed the phylogenetic development of each tumor, as described previously.\textsuperscript{5} Details about this approach are provided in the Supplementary Methods section and in Figures S1 through S3 in Supplementary Appendix 1; an
overview of the two cases is provided in Table S1 in Supplementary Appendix 2, along with details about the two patients in Tables S2 through S5. The two appendixes are available with the full text of this article at NEJM.org.

**PATIENT 1**

In Patient 1, we examined inherited and de novo germline variants. We found two bona fide pathogenic mutations, one inherited from each parent. The child carried a paternal truncating variant in *SMARCA4*, which is a putative predisposition gene for neuroblastoma.\(^7\)\(^9\) In addition, he carried a maternally inherited truncating mutation in the gene encoding checkpoint kinase 2 (*CHEK2*), which has recently been found to confer a predisposition to a variety of childhood tumors\(^7\)\(^9\) (Fig. 2B). Next, we examined mosaic variants that were present in one or both tumors and in blood cells. These variants are nonheterozygous in the blood and can be inferred to have accumulated during the first few cell divisions of the zygote. Three of these variants were present in blood and in both tumors (Fig. 2C). Unexpectedly, four of these variants were present in the left tumor and blood but absent in the right tumor. These variants, which are exclusive to cells of the blood and the left tumor, were not found at chromosomal loci affected by the loss of heterozygosity or copy-number variation at chromosomal loci (Table S5 in Supplementary Appendix 2). The exclusivity of these variants indicated that the tumors had been derived from separate embryologic lineages that segregated within the first few cell divisions before gastrulation. In a separate analysis, we did not find any somatic variants that were shared between the left and right tumors, which was consistent with the previous finding of exclusivity.

After segregation of the two lineages, the tumors had undergone remarkably parallel development (Fig. 2D). Among key somatic changes in each tumor was the gain of chromosome 7, albeit of different parental chromosomes: duplication of the maternal chromosome in the left tumor and duplication of the paternal chromosome in the right. Furthermore, both tumors had a loss of maternal 19p, thus deleting the remaining intact germline copy of *SMARCA4*. However, different translocations generated the breakpoint on 19p (Fig. 2D, 2E, and 2F). In the right tumor, the loss of 19p was coupled with a gain of 17q, a genomic feature of high-risk neuroblastoma.\(^6\) In addition, on chromosome 11, the right tumor had a high density of intrachromosomal rearrangements associated with copy-number changes, which was consistent with chromothripsis (i.e., multiple chromosomal rearrangements that are thought to occur simultaneously and are commonly found in human cancer). To establish the order of these parallel somatic events, we applied analytic techniques that time the acquisition of copy-number changes relative to that of single-base variants.\(^4\)\(^10\) Our findings indicated that these somatic events may have been acquired in the same order on both the left and right sides (Fig. 2E). As has been shown in other cancers,\(^4\)\(^10\) we identified trisomy 7 as an early acquired event. Among the somatic events unique to each tumor, we did not find any additional driver events.
Patient 2

We performed the same analyses on samples obtained from Patient 2 and her parents, and we again found 3 variants in both tumors and in blood, along with 14 variants that were exclusive to one tumor or the other yet present in blood cells (Fig. 3B and 3C and Fig. S4 in Supplementary Appendix 1). We did not find variants that were common to both tumors in a separate analysis. We concluded that both tumors were derived from independent embryonic lineages that segregated within a few cell divisions after fertilization. In Patient 2, tumor development diverged, leading to a low-risk neuroblastoma in the right adrenal with no additional driver events. By contrast, the tumor in the left adrenal had acquired somatic genetic changes that characterize high-risk tumors, including gain of chromosome 17q (Fig. 3D and 3E), consistent with its characteristics (locally and extensively invasive).

Discussion

Here, we reconstructed the development of two cases of bilateral neuroblastoma from somatic mutations that provide a barcode for embryonic cell divisions and lineages. Our expectation might have been that such tumors stem from a common postzygotic precursor, similar to what has been observed in bilateral Wilms’ tumor. In such a case, we would have observed shared mutations between tumors that we did not find in blood. However, we found that such tumors...
are independent neoplasms that separated early in embryogenesis, thus providing direct phyloge-netic proof for the presumption that some multi-site neuroblastomas do not represent metastatic disease.

Although our study is limited in scale owing to the rarity of bilateral neuroblastoma, our findings raise questions about multisite tumors in children. In the majority of cases, it is likely that, as commonly assumed, multisite tumors represent metastatic disease. Nevertheless, our observations may lead to some questioning of this assumption in certain clinical scenarios, such as multisite tumors in infants or in children who are predisposed to cancer. One could argue that for a child with independent multisite tumors, treatment should be guided by the lesion associated with the worst prognosis, with a less aggressive treatment regimen in the instance of multiple, independent low-risk tumors.

In our study, Patient 1 had two embryonic lineages that evolved in parallel into high-risk tumors. Although the parallel convergent development of different clones within a tumor is a well-described phenomenon,\textsuperscript{11} the development of independent synchronous tumors that mirror each other is a remarkable observation. We speculate that each of the germline pathogenic variants sets the stage for tumor formation through a restricted repertoire of additional driver events. Our analytic strategy may provide further insights into the early embryonic development of childhood cancers, especially because whole-genome sequencing in the context of clinical management of cancer is becoming increasingly common.

Supported by Children with Cancer UK and Wellcome.

Disclosure forms provided by the authors are available with the full text of this article at NEJM.org.

We thank the children and their families for participating in our research.

REFERENCES